PAGE NO.	1	going !
DATE:	1	1

Introduction to Medicinal Chemistry

Medicinal chemistry as a subject explains the design and production of come organic compounds that can be used for the prevention, treatment or cure of diseases.

According to Burger medicinal chemistry tries to be based on the ever-increasing hope that biochemical valionales for drug discovery may be found." In practice medicinal chemistry is based on the hope of discovering biochemical pathways as well as modification of structurer come having known physicathemical physiologic or pharmacologic effects.

The primary function of medicinal chemist is still to discover new angs and the knowledge of principles of brochemical action are proving to be very helpful for design of new drag molecules.

In the ancient period, natural products having history as folk medicine were used for drigg therapy but now a days very little of these remedies are used. The molecular orbital and other calculations that elucidate the electronic and conformational aspectation of molecules are molecules are more used its predict the

PAGE NO. 2

shuchires for delective biological

for therapy began in early nineteenth with the use of chloroform & ether anesthesia. The late nineteenth century dominated by Paul Ehrlich. This period tremendous development in medicinal

Lew a days the drug discovery process is a cam work which includes scientists from various disciplines including biology, toxicology, pharma-cology, mison biology & bropharmary.

Figure 1.1 Contributors to medicinal chemistry.

Pharma cognosy

Analytical Chemistry

X-ray coxetallopaphy

Spectroscopy

Spectroscopy

Chemistry

CHEMISTRY

Microbiology

Microbiology

Analytical Chemistry

Spectroscopy

Medicular Pharmacology

Physical Chemistry Computational Chemistry

Microbiology, Immundogy & Genomics

Reference
M.E. Wolff, Burger's Medicinal Chemistry, 5th ed., Ret
Part I, The Bouris of Medicinal Chemistry, Hew York,
Wiley-Interscience,

Chapter 2



Physico-chemical Properties and & Drug Activity.

Physico-chemical properties refer to the influence of the organic functional groups present within a molecule on its acid/base properties, water solubility, crystal structure, partition coefficient, etc. In design of better medicinal agents the relative contributions of each functional group adds to the overall physical and chemical properties of the motecule.

. Selectivity of Drug Action at Active Site

Ehrlich gave the concept of drug receptor. It stated that certain "side chains" on the surface of cell were "complementary" to the drugs and hence allow the two substances to combine.

the st relectivity of doing action was rhown via the concept of "magic bullet" for compounds that diminish the disease states without producing unwanted harm to the organism being treated. The structural elements (functional gradue) within a molecule contribute in an additive manner to the physico-chemical properties of a molecule and hence to its biological action.

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Physico-chemical Properties of Drug Molecules.

The most pharmacologically influencial physico-chemical properties include:

(a) Acid-Base properties

(b) Relative Acid Strength Cpkal (c) Water Solubility

Acid-Base Properties when considering the solution behaviour of drug within the body we deal with dilute solutions. Lowry-Bronsted acid-base theory explains and predicts the acid/base behaviour

of doing molecules. The acid-base properties of doing molecules directly affects absorption excretion and compatibility of drug with other drugs in solution. According to Bronsted theory, and is any substance expable of donating proton [H+] in solution whereas base is any substance capable of accepting proton (Hi) in solution. The acid gives a proton to base and is thus converted into its conjugate base.

Example! CH3COOH + H2O = CH3COOF + H3OF (Acid) (Base) Conjugate Conjugat
Base Acid

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The base on the other hand, accepts a proton from and acid & is converted into its conjugate acid.

Example: CH3NH2 + H2O = CH3NH3 + OHD

[Base] (Acid) (anjugate (anjugate Azid Base)

when an acid loses its proton, it has extra pair of electrons which do not neutralize by the proton. This is the ionized form of an acid and is highly water soluble & due to change. The acid is said to the have undergone dissociation.

1. Phenol REOTOH Phenolate REOTO

2. Alkyl thiol R-SH Thiolate R-SE

when a base is converted to its conjugate acid, it is also ionized and carrier a positive charge due to extra proton. Most basic drugs are usually derivatives of primary, secondary and tertiary amines.

Base Conjugate Acid Anyl Ammonium REOTHY.

ate 1. Anyl Amine REOTHY.

2. Imine R-C=N-H Iminium R-C=NH2

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Organic functional groups that neither give up a proton nor accept a proton are said to be neutral with respect to acid-base properti-Example ::

R-CH2OH; R-O-R; R-C=H Alkyl Alcohol Ether nitrile

A molecule may contain multiple functional groups and therefore possess both acidic and basic properties. For example; Ciprofloxacin. It contains secondary amine and a carboxylic acid group. Depending on pH of the solution, the molecule can either accept or denate a proton or both. Thus it can be acidic, basic or

amphoteric o neutral COOH] acidic weak base

At a given pH value only one functional group is ionize

Relative Acid Strength Cpka]
The concept of pka indicater the relative acid / base strength of organic functional groups and allows to calculate, for a given pH, the amount of molecule in the ionized and unionized form.

Strong acids and bases dissociate or accept proton to produce their respective conjugate bases and acids.

 $HadH + H_2O = Ha^{\oplus} + OH^{\oplus} + H_2O$ $Hcl + H_2O = (l^{\oplus} + H_3O^{\oplus})$

water is amphotenic. In dilute agreeous solution the strongest base present is OHE and the strongest acid is H_3OD . This is called as levelling effect of water.

Predicting the degree of ionization of a molecule. To predict the degree of ionization of any molecule, the pla values of the acidic and basic functional groups present in the molecule should be known. Handerson-Hasselbach equation is used to calculate the percent ionization of compound at a given pH.

pka = pH + log [acid form]
[base form]

3

ON

en

the	percent	ionization	of a drug	is calculated
using.	Eq. 1-1.	for HA	acids Chonto	inized) and
· Eq. 1-9	2 for	BH+ acids	(ionized).	is calculated and
				^

% ionization =
$$\frac{100}{1 + 100(Pka-PH)}$$
 (Eq. 1-1)

Table 1-1 gives an index of the effect of PH& pha on the percentage ionization of HEA HA acids and BHT acids.

1able 1-1	r		. 4
Percentage	lonization	Relative to pka	
		lonization(%)	

	HA Acids	BH+ Acids	
pka-2 pH units	0.99	99.0	
pka-1 pH unit	9.1	90.9	
pha = pH	50.0	0.02	
Pka + 1 pH unit	90.9	9.1	
Pha +2 pH units	99.0	0.99	

pha affects the distribution of drug molecule in various tissues of the body.

Water Solubility of Drugs
The solubility of drug molecule in & water affects the routes of administration, absorption, distribution and elimination.
The hydrogen bonding potential in the molecule and the ionization of functional groups are considered in study of water solubility of molecules.

Each functional group capable of donating or accepting a hydrogen bond contributes top overall water solubility of the compound. Such functional groups increase the hydrophilic nature of molecule. When two molecules containing dipoles approach one another, they align such that the negative end of one dipole is electrostatically attracted to the positive end of the other. If the positive end of dipole is Hydrogen adom, it is called as hydrogen bonding. for a Hydrogen bond to acce occur at least one dipole must contain an electropositive hydrogen atom.

Several possible H-bond types may occur with different organic functional groups and water. different organic functional groups and water.

PAGE NO. (O REDUCE)

More the hydrogen bonds, the higher is the water solubility.	(
woder soldbility.	. 1
Table 1-2 exemplifier certain function groups and their potential no or number of H-bonds.	n
and their potential no or number of H-bonds.	H
Table 1-2	
Functional Groups and H-bonding	
	T
Potential H-bonds	11
R-OH	
$R-NH_2$ 3	TI
R-C-R'	
	C
R-N-H 2	M
	P'
R'	60
R-M-R'	av
R"	For
lonization	
lon-dipole bonding plays an important vole	(,
in determining the water solubility of molecule.	
ion-dipole bonds develop between a carion	M
lon-dipole bonds develop between a cation or anion and a formal dipole like water.	
A cution associates with negative end of the	and the same of th
CIPOLE.	
An anion associates with positive end of the	
CLIDOLE.	t
In ion-dipole bonds, as in organic salts, to	

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associate with enough water molecules to become water soluble, the salt must be highly dissociable.

Highly dissociable salts are formed from

(a) strong acid and strong base

(b) weak acid and strong base

(c) strong acid and weak base

The strongest acids are HCL, HNO3. H2SO., , perchloric and performic acid. The strongest bases are NaOH and ROH.

Compounds with ionizable functional groups that produce opposite charges can interact with each other rather than water. Such compounds are woder insoluble. For Example; Amino acid Tyrosine

Greater the separation between charges, higher the water solubility of molecule.

Predicting water Solubility

the carbon solubilizing potential of the functional groups.

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More the hydrogen bonds, the higher is the	a
woder solubility.	pe
Table 1-2 exemplifier certain function groups	hi
Table 1-2 exemplifier certain function groups and their potential no. of number of H-bonds.	H
Table 1-2	
Functional Groups and H-bonding	
Function Group Number of Potential H-bonds	Th
R-0H 3	r
$R-NH_2$ 3	Th
R-C-R'	•
	Co
R-N-H 2	pr
R'	ea
R-M-R'	Q.V(
R"	For
lonization	
lon-dipole bonding plays an important role	
in determining the water solubility of molecule.	9
lon-dipole bonding plays an important role in determining the water solubility of molecule. Ion-dipole bonds develop between a cation or anion and a formal dipole like water. A cation associates with negative end of the lipole.	hic
or anion and a formal dipole like water.	- (
A cution associates with negative end of the	
An anion associates with positive end of the	
dipole.	H
dipole. In ion-dipole bonds, as in organic salts, to	F

If the solubilizing potential is more than the total number of Carbon atoms in the molecule, the molecule will be water soluble. Functional groups that can form intramolecular H-bonds decrease the solubilizing potential and there fore decrease the water solubility.

2. Analytical Approach !- It involves the calculation of approximate Log P. value or the log of the partition wefficient of the molecule.

Partition Coefficient.

It is the vario of the concentration of day in octanol to that in water.

Log P is a measure of the solubility characteristics of molecule.

A hydrophobic / hydrophilic value [hydrophobic substituent constant; TT] is given to each functional group.

Log P = ZT

Water solubility is the solubility of more than \$3.3%; a equivalent to about 0.5 Log P.

Log P values less than +0.5 tend to increase water solubility and Log P values more than +0.5 tend to decrease water solubility.

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lonization state of a molecule influences its water solubility and the ability to of the molecule to traverse biological membranes and hence its ability to get absorbed in body.

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rel

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Stevenchemical Features and Pharmacological Activity

The physicochemical properties of a drug molecule are dependent upon the functional groups present in the molecule and their spatial arrangement (steres chemistry). A drug molecule is subjected to many complex processes from its administration to elicitation of biological response (fig. 3-1)

Figure 3-1 Drug Administration to Biological Response

Stereo chemistry of the molecules play a major role in the pharmacological properties, as many of these processes are stereospecific. Stereo chemistry is responsible for the difference in degree of pharmacological activity of isomers. The structural features required

The influence of steric factors on pharma-

cot cological action is categorized in three groups -(1) Optical and geometric isomerism (2) Conformational isomerism

(3) bosterium & pharmacological action.

Optical Gomerian

Optical le Geometric Lomerism And Pharmacological Activity

Optical Isomerism
Optical isomers are compounds that differ only in their ability to notate the plane of polarized light. Optical isomers may exhibit different biological activities. for example, one isomer of ammonium tartarate inhibits the growth of Penicillium glaucum whereas the other isomer has no effect.

Optical isomers may be enantioners or diaster -eomers.

Enantiomorphy are non-superimposable mirror images. They rotate the plane of polarized light in equal amounts but in opposite direction

Diastereomers are isomers that are neither mirror images nor superimposable.

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Influence of optical bornerism on Pharmacological Activity. The differences in biologic activity between optical isomers depends on their ability to react selectively at an asymmetric center in the biological system. The Figure 3-2 exemplifies the affect of these difference. Fig. 3-2 Effect of Garneric Structure \$ biologic system of the two isomers shown in above figure, only one (I) has the correct orientation for all the three groups to fit at their respective sites on the biological system (receptor) and therefore only (I) is active biologically. This is called as Easson - Stedman Hypothesis or the three point fit hypothesis. The differences in distribution of the isomers in the biologic system may also lead to the

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differences in pharmacological activities. The ifferences in distribution occurs because the somes may be selected by some other asymmetric center in the system before it reaches the specific receptor. Figure 3-3 its subjected to before reaching the specific receptor.

Fig. 3-3 Selective processes in Drug Action.

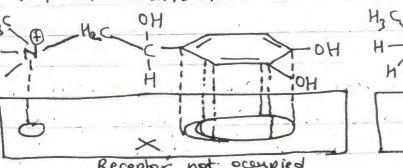
Drug Dose > Membrane > Selective Metabolism

Drug Dose > Membrane > Selective Metabolism

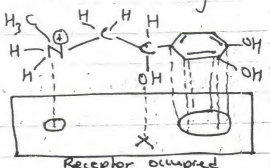
Drug Dose > Membrane > Selective Metabolism

Desired Eng Hon specific Response Receptor receptor. [Site of Loss]

For example, only the (-) isomer of epinephine has the -OH group in correct orientation to allow perfect binding with all groups to the receptor. Hence (-) epinephine has high pressor activity whereas the (+) epinephine is last in dishibution & has minimal activity.



Receptor not occupied t) Epinephine leu autivity



(-) Epinephine - Mare Autive

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steven chemistry plays role in the metabolism of optically active drug molecules. On binding with vacemic drugs the metabolizing enzymes produce diastereometric complexes and therefore lead to different rates of metabolism: stereo-selective meatabolism. A metabolized drug may have increased or decreased activity. But example L-(+) lackly choline hydrolyzes much readily than D-(-) lackly choline.

Selectivity of passage of drug through membrane occurs due to asymmetric centers within the membrane. If the drug has to cross the membrane

Selectivity of passage of drug through membrane occurs due to asymmetric centers within the membrane. If the drug has to cross the membrane to reach its receptor then selectivity at the membrane is important in biological activity. The transfer of molecules across a membrane by Permeases is a selective process. For example, only L-isomer of valine, lewine penetrate the cell wall of bacteria such as E.coli whereas the D-isomers do not.

Geometric Isomerism (cis-trans isomerism)

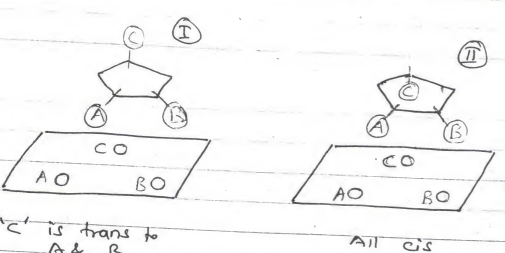
It indicates a type of diasteremen that focus as a result of restricted notation around a bond.

Then, certain identified groups are present on is some side of the plane of molecule, the difference molecule is said to be cis.

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when the identified groups are on opposite sides of the plane, then the molecule is said to be trans be trans

Influence of Geometric Isomerism on Biologic Adivity The effect of geometric bornerism at the receptor site is shown in figure 3-4 fig. 3-4: Geometric isomers on receptor site



Three substituents of cyclopentane sing (A,B&C) are needed for binding to the receptor surface. Only the 'cis' arrangement (II) allows

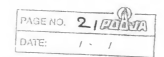
this and hence if gives biologic activity. or example, trans-Diethylshilbesterol is 14 times where than the cis-isomer. he differences in biologic activity of geometric omers may be due to differences in interatomic istance of the groups essential for pharmacologic response.

Conformational Isomerism And Biologic Activity
Conformational isomerism is defined as
the non-identical spatial arrangement of
atoms in a molecule, regulating from
rotation about one or more single bonds.
for example, Hydrogen Peroxide (4202) gives
distinct conformations on rotation about the
D-H bond.

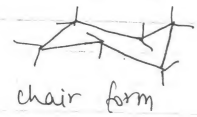
H-0/0 H-0/H

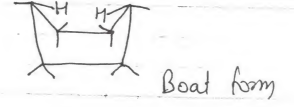
Ong molecules are complex structures. The barrier to free rotation about single bonds in drugs is due to the decreasing distance between the H-atoms on the adjacent carbon atoms on the C-C bond is rotated.

For example, Ein eclipsed conformation, the hydrogen atoms are in closest proximity and hence the molecule is unstable. The staggered conformation, on the other hand, gives allow the greatest separation of hydrogen atoms and hence is the most stable conformation.



Cyclo hexane exists in two conformations: hour and boat. The chair conformation spreferred over boat because in chair onformation all the bonds are staggered.





substituted rigid molecules, the axial substients are in entirely different environment from unitorial substituents and therefore differences in systeal or chemical properties.

Influence of Conformational Isomerism on Biologic Activity

pon interaction with substrate 1; the enzymer indergoes conformational change. Similarly

on interaction with day molecules, the receptor ordengoes conformational change.

receptor site may bind to only one of the any conformations of a day molecule. The olecules that can adapt the conformation eded for binding for binding may act as oniat or autoconist. onist or antagonists.

Antagonists bird to receptor but do not elicit ponse due to lack of some groups.

For example, groups A&B are needed for binding to receptor and C is needed for response (Figure 3-5)

Figure 3-5:

H FA A

H B B

Concept of Agonist & Antagonist

(C)

HA-I-A

HA-I-A

HA-I-B

HA-I-B

I. Agonist

II. Antagonist No 'c' group; which is needed for response

MI. Antagonist & optical bromen of I. Can bind but no response

Conformational Formers analysis explains the differences in biologic activity of diastereometric drugs.

Isosterisms and Pharmacological Action
Isosterism relates to the similarity in physicochemical properties of atoms, grups radicals and molecules with similar electronic phractures.

Grimm's concept of hydride displacement vertical columns of isosteric groups are formed by displacing one place to the right successively the elements of a row & adding a hydrogen atom. Molecules of an isosteric pair should fit in the same crystal lattice.

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Table 3-1 \$

Iso	steric	Pairs	of Grimm's	Concept
C	11	0	F	Me
	CH	MH	ОН	HF
		. CH2	NH2	OH2
			CH3	NH3
				CH4

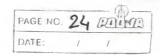
Each vertical column represents a group of isosteres.

Bio isosknism

Application of the concept of isosterism to modify the biological activity is called as bio isosterism.

Classification of Bioisoskres

- I. Classical Bioisostères
- (1) Monovalent atoms or groups. Example, halogens & -XHn, X is C, N, O & S
- (2) Divalent atoms or groups. Example, R-O-R': R-HH-R'
- (3) Trivalent atoms or groups. Example, R-H=R'; R-CH=R'
- (4) Tetra substituted atoms. Geample, = C=;
- =H=; =P=
- (5) Ring Equivalents. Example, -CH=CH-; -S-; -O-; -NH; -CH2



Hon-classical Bioisosteres

(1) Cyclic us nonexolic bioisosteres Example, prome thatine -> methodilazine (2) Exchangeable Groups

(a) Hydroxyl group bio isos keres

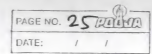
(b) Carbonyl group bio isos keres

(c) carboxylate group bio isos keres

(d) Amide group bio isos keres

(e) Thio urea biosos keres

(f) Halogen bio isos keres



Drug Receptor Interactions lost of the pharmacologically active agents re structurally specific drug molecules. pear to be having greater influence on drug effect. A direct interaction of drug th receptor material initiates a sequence events leading to response. receptor is defined as a tissue reponent which fulfils the following itéria it is a macromolecule which has to having chemorecognitive properties for a ecific dangs; the specificity for sites on receptor and the inction of the receptor are genetically determined binding of agonists [endogenous wastance/ ug] to initiates a chain of events leading response; and the binding of agonist at the receptor e does not depend on any bond making or eaking in the agonist molecule.

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	Th
The quantitative ability of a drug do interact with the receptor is called as affinity.	40
with the receptor is called as affinity.	Cle
The ability of the drug once bound to the receptor, to produce its biological response is	· Conc
receptor, to produce its biological response is	er.
alled as its expose inthinsic activity on	die
efficacy.	hage
	Che
Types of Drug-ReceptorfoxInteractions.	6-11
9	incl
The interaction of many structurally specific negs with receptor is as under. $D+R = \frac{1}{k_2} DR - \frac{1}{k_3} E Eq. 4-2$	For
nigs with receptor is as under.	inte
$D+R = \sum_{k_2} DR - \sum_{k_3} E \qquad (q. 4-2)$	Ech
	with
he two step sequence involves an equilibrium	anic
etween the doing and neceptor. Both steps are	Covo
trongly influenced by drug-receptor bonding & se by stereochemical fit of the drug on he receptor. The forces involved in the bonding	eft
by stereochemical hit of the drug on	
he receiptor. The forces involved in the banding	10
f drug and receptor include covalent bonds,	The
nic, re-inforced ionic hydrogen bonds, ion-dipole, ipole-dipole [keesom] forces Van der waals	inte
upole-dipole [keeson) forces Van der waals	(1)
London forcer Cinduced dipole-induced dipole) and	The
leybe's forcer (dipole-induced dipole) and the	is
ydrophobic interactions.	opp
	5
Covalent Bonds	9-(
Mutual sharing of electron pairs between up atomosphis produces a covatent band.	d-(
us alaman produces a covoitent bond.	a

The bond strength of a covalent band is 40-140 kcal/mol. Such bands do not cleave spontaneously under physiological conditions. Cleavage occurs under ensymptic or specific acid-base catalysis. The effect of drug terminates once the drug-receptor band has cleaved.

Chemical mechanism that lead to covalent band formation between drug and receptor include alkylation, acylation & phosphorylation. For example, the reactive immonium-ion intermediate of anticancer nitrogen mustards [chloram bucil] readily forms (ovalent bands with sulfhydryl, carboxylate and phosphate anions and with unchanged MS & O atoms. Covalent bands may produce irreversible effects.

Mon-(ovalent Bonds
These bonds produce short-lived & reversible interactions. They have low bond strengths.

(1) Ionic Bond.

Thise are is formed by transfer of electrons. It is the electrostatic attraction between the oppositely charged ions. The bond strength is 5 kcat/mol. Functional groups Juch as a carboxy [[terminal], secondary phosphory], a - ammonium terminal present on receptors; and aliphatic amino, quaternary ammonium

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groups present in drugs are ionized at the physiological pH and need lead to the formation, of ionic bonds. iv 81 (2) Hydrogen Bonds It is a strong dipole-dipole interaction in which hydrogen atoms serve as a bridge between two electronegative atoms, holding one by covalent bond and other by pure electrostatic forces. X-H----Y to this bond 'H' is covalently bonded to'x' & ionically to 'T'. The positive pole of one dipole is hydrogen atom (x=Hs+). The most frequent hydrogen bonds occur between -oH group and the NH groups in the following order of decreasing stability.

OHH > OHO > NHH = NHO 3 Hydrogen bonds may be intermolecular or inframoleular. In aqueous medium, all free H-bonding groups on drug and receptors are linked to water through H-bonds. The bond strength of H-bond is 1 to 7 healprol.

Formation of H-bond leads to agreens solubility
of drug which is necessary for drug

molecules to be transported to the site of

action on a receptor

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These are short range forces involving bonding interactions with non-polar groups. The bond strength may be 0.5 to 1.0 kal/mol. In the medicinal agents, non-polar groups help in strengthening the drug-receptor interactions & in assuring the appropriate water-lipid solubility relationship.

London forces lead to bonding between two non-polar groups and are created from induced dipoles which arise from polarization of electron clouds.

When the non-polar moiety is of sufficient size and of appropriate steric configurations London forces may stabilize a duag-receptor complex.

These are the most important interactions involved in maintaining the normal configuration of proteins and in determining the biological effect of many drugs. The bond strength of hydrophobic interactions is theal/mol. The interaction energy is related to gain in entropy for the system.

Each water molecule is H-bonded & Fair other with four bonds to neighbouring water molecules is highly ordered.

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dipole-induced dipole bonding interactions is extremely weak and the energy gained is not enough to compensate for the increased order which results from dispersion of solute in water.

The non-polar solutes have low agreems solubility and tend to aggregate in agreems solution, freeing the ordered water molecule a thus increasing the entropy of the system. The complementary fit of a doing on reception may require a close approach of non-polar residues, which can be facilitated only by freeing of water molecules between the two besidues.

Hydrophobic bonding constant, $\pi = \text{Log}P_x - \text{Log}P_H$ where $P_x \& P_H$ are partition welficients of
substituted and parent compounds respectively.
Postive π values increasing hydrophobic bording
whereas regative π values decrease hydrophobic
bonding

Stereochemistry of D-R Interactions.

An exact fit of the drug molecule and receptor is necessary for maximum response: Most of the structurally specific drugs act stereospecifically when they exist as configurational isomers.

Theory of D-R Interactions Receptor theory involves the ensyme kinetic

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model based law of mass action $K_d = \frac{[D][R]}{[DR]}$ [OR] The response of drug is recentary dependent on the per number of receptors on a given tissue and the affinity of the receptor for drug. Apprists bind to the receptor and lead to activation intracellular components involved in the physiological responsiveness of the cell or tissue. Antagonists bind to the receptor and block the interaction of agonist. They do not produce effect of their man Inverse agonists interact with a defined recognition site on the receptor and are not only able to block the effects of agonist but are also able to produce their own effect opposite to the equivit. five theories have been proposed for D-R interaction) Occupancy theory (2) Rate theory (3) Inactivation reory (4) Induced - Fit theory (5) Macro molecular exhapsion theory.

Occupancy Theory
The basis of occupancy theory is that the effect
produced by an agonist is dependent on the
number of receptors occupied by the agonist.
Dring the Michalis-Menkn derivation of law of
made action, the occupancy theory states that-

(i) the D-R complex is reversible;
(ii) the association of dong with receptor to form D-R complex is a proctor bismolecular process while dissociation is unimolecular;
(iii) all receptors of a class are equivalent as bind to the dong independent of each other;
(iv) formation of D-R complex obesnot after the affinity of the receptor for the dong;
(v) the response is directly proportional to the number of receptors occupied; and
(vi) the biological response is dependent on the attainment of equilibrium between the drug receptor.

The interaction of antagonist with the receptor results in occupancy without elicitar of a functional response.

Rate Theory.

According to the rate theory, the respective to an agonist day depends on the rate of D-R complex formation. The effect 'E' is given as E = 0 Veg E = 0 Veg

The rate of D-R complex formation changes the receptor mediated events. The rate of association or dissociation of agonist is rapid and leads to sequence of impulses. Antagonist has high association constant to

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a low rate of dissociation.

Inactivation Theory
This theory is a hybrid of both occupancy and rate theory. This theory assumes that the D-R complex is an intermediate 'active' state that gives rise to an inactive form of the receptor, R' that is part of an D-R complex (R'-D) ky is rate of association of D-R is rate of association of D-R is rate of dissociation of D-R is rate of dissociation of D-R to R'-D at the rate constant for regeneration of R from 2'-D is ky 21-D is k3 $[D] + [R] \xrightarrow{k_{+1}} [D-R]$ [R'D] " low the response of drug is proportional to he rate of R' formation which is k3[R'-D]. I depends on both the rate of formation of and the number of receptors occupied.

Induced fit Theory
he occupancy a rate theories do not provide
pecific models at the molecular level to
count for drugs acting as aganist or antagonist.
Induced fit theory is based on induced—
t model of enzyme—substrate interaction leading
conformational change in enzyme and hence

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active orientation of groups.

It assumes that protein constituents of the biologic membrane play a note in regulating ion flow.

The drug [for eg. Acetylcholine] may interact with the protein and after the normal forces that stabilize the structure of the protein and hence produce a transient change rearrangement in the membrane structure and the consequent change in its ion-regulating properties.

Macromolecular Perhirbation Theory
This theory is explained with the help of mode of action of acetylcholine at the musicannic receptor
Interaction of small molecules [drug] with a macromolecule [Receptor] may lead either to Specific Conformational Perhirbations [Scp] or to non-specific Conformational Perhirbations [NSCP] A SCP results in the specific response of an agonist.

If an NSCP occurs, an antagonistic action make produced.

If a drug possesses features that contribute to formation of both SCP and NSCP it results in partial stimulation action [partial agonist].

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a low rate of dissociation.

Inactivation Theory
This theory is a hybrid of both occupancy and rate theory. This theory assumes that the D-R complex is an intermediate 'active' state that gives rise to an inactive form of the receptor, R' that is part of an D-R complex (R'-D) ky, is rate of association of D-R ky, is rate of association of D-R ky, is rate of dissociation of D-R to R'-D ky, is rate of dissociation of D-R to R'-D ky, is rate of dissociation of D-R to R'-D ky, is rate constant for regeneration of R from 2'-D is ky. $[D] + [R] \xrightarrow{k_{+1}} [D-R]$ E9\$ 4-5 [R'D] K low the response of drug is proportional to re rate of R' formation which is k3[R'-D].

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active orientation of groups.

It assumes that protein constituents of the biologic membrane play a note in regulating ion flow.

The drug [for eg. Acetylcholine] may interact with the protein and after the normal forces that stabilize the structure of the protein and hence produce a transient change rearrangement in the membrane structure and the consequent change in its ion-regulating properties.

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If an NSCP occurs, an antagonistic action must be produced.

If a doing possesses features that contribute to formation of both SCP and NSCP it results in partial stimulation action [partial agonist].

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Figure 4-1 Specific & Hon specific Conformational Perturbations. SCP [Agonist Action] SH Me3 Cg MMe3 NSCP Antagonistic State mysigninic receptor SCP (Shimwant Action) MSCP [Blocking Action] Partial Agonistic Action.

DATE: / /

Prodrugs

Almost all dougs possess some underivable physicochemical properties and biological properties while designing a new doug delivery system three factors must be considered—

(i) doug component; (ii) vehicle / carrier; and (iii) intended route of administration.

Drug Component.

1) Hard Drug: It is resistant to biotransformation and has a doing biological half life. Design of a hard drug inholves the metabolic stabilization of existing molecules by replacing functional groups susceptible to to biotransformation by sloble grow This is also to called as derivatization.

For example: Stabilization of tol-butamide by replacing CH, by CL as in chlorproparnide.

13C SoznHC-NHC4Hg CL OSZNHC-NHC4Hg

Tolbutamide

Chlorproparnide

(2) Soft Dong: It is a biologically active compound which is biotransformed invivo in a rapid and predictable manner into non-toxic metabolites.

Design of soft dongs involves the concept of metabolic states witching in which a functional